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RESERVE

PATENT SPECIFICATION

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COMPLETE SPECIFICATION

Dibenzoate Esters of Aliphatic Diols and process for their preparation

We, UNION CARRIDE CORPORATION, (formerly known as Union Carbide and Carbon Corporation), of 30, Bast 42nd Street, New York, State of New York, United States of America, a corporation organised under the laws of the State of New York, United States of America, (assignee of Gordon Murray Goodale, Edward James Mills, Jr., Thomas Robert Miller and James John 10 Fassiacht), do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

This invention relates to a method of pre-

This invention relates to a method of preparing dibenzoate esters of aliphatic diols by reacting buryl benzoate with an aliphatic diol containing four to nine carbon atoms and having no tertiary hydroxyl groups. A tertiary 20 hydroxyl group is one which is attached to a

tertiary carbon arom.

Among the compounds which may be prepared in accordance with the present invention are the dibenzoate esters of aliphatic diols having an oxygen interrupted carbon chain, such as diethylene glycol dibenzoate, dipropylene glycol dibenzoates and the dibenzoate esters of straight and branched chain aliphatic diols having an uninterrupted carbon chain, such as 2-ethyl-1,3-pentanediol; 2-methoxymethyl-2,4-dimethyl-1,5-pentanediol; 2-ethyl-2-butyl-1,3-propanediol; 2-methyl-1,3-propanediol; 2-methyl-1,3-propanediol; 2-methyl-1,5-pentanediol; 2-ethyl-1,5-pentanediol; 2-ethyl-1,5-pentanediol; 2-ethyl-1,5-pentanediol; 2-ethyl-1,5-pentanediol; 2-ethyl-1,5-pentanediol; 2-ethyl-1,5-pentanediol.

Dibencette at 1,5-pentanediol.

Dibenzoate esters of aliphatic diols of the 40 class described above are useful as plasticizers for vinyl resins. Dipropylene glycol dibenzoate is particularly valuable as a plasticizer for polyvinyl chloride resins.

We have discovered that the dibenzoate esters of aliphatic diols which contain from 4

to 9 carbon atoms and which containing no tertiary hydroxyl group can be prepared by reacting one molecular proportion of one of the above described aliphatic diels with 2.5 molecular proportions of buryl benzoate in the presence of an alkaline catalyst. The reaction to be conducted at a temperature of from 100° C. to 250° C, and preferably from 150° C to 180° C, for a period of from 10 hours to 30 hours. Although the reaction can be conducted at a pressure of from 0.1 mm. Hg to 200 mm. Hg, it is preferably conducted at a pressure of from 2 mm. Hg to 100 mm. Hg. Snimble catalysts for the reaction compaise alkali metal and alkaline earth metal oxides, hydroxides, alkoxides, carbonates and borates. Preferred catalysts are the alkaline earth metal oxides. Calcium oxide is a particularly suitable catalyst because of its good catalytic activity, low cost and ease of handling. Catalyst conlow cost and ease or handling. Cararyst con-centrations can vary from 0.06 per cent by weight to 1.4 per cent by weight but are pre-ferably between 0.1 per cent by weight and 1.0 per cent by weight. By operating at a tem-perature in the range of from 158° C. to 180° C. with a catalyst concentration of 0.1 per cent by weight, excellent results are obtained, particularly in the production of dipropylene glycol dibenzoste. As the reaction proceeds, burned is removed by distillation, followed by a mid-fraction, usually containing butanol and butyl benzoate and a third fraction of butyl benzoate. The pure dibenzoate ester of the aliphatic diol can be isolated from the residue by distillation or by filtration.

The following examples are illustrative:-

Example I.

Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Raschig rings and surmounted by a standard reflux head with the usual condenser and vacuum receiver attached, were charged 8015 grams (45.0 mols) of refined buryl benzoate and 2414 grams (18.0

B.

120

mols) of dipropylene glycol. Then 104 grams (1.0 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a pressure of 100 mm. Hg. Buranol was removed by distillation at this pressure until the kettle tem-perature increased to 180-194° C. The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg, and the 10 remaining butanol was collected as distillate. A mid-fraction, containing butanol and butyl benzoate was then distilled, followed by a pure fraction of butyl benzoste. Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature below 198° C. The contents of the reaction kettle were cooled, mixed with a filter aid and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a goose-neck head and an external nitrogen ebullans. The stripping operation was conducted at a temperature of 182° C at less than 1.0 mm. Hg. The kettle residue was then 1.0 mm. Hg. The kettle residue was then treated with a decolorizing material, filtered, and recovered.

A yield of 81.7 per cent of dipropylene glycol dibenzoate (based on dipropylene glycol) was obtained, with an efficiency (based on butyl benzoare) of 86.4 per cent. The 30 refined dipropylene glycol dibenzoare had a purity by saponification of 99.8 per cent, an acidity (as benzoic acid) of 0.02 per cent, a diol content of 0.69 per cent and the following physical properties:-

Refractive index at 20° C. Specific gravity (20/20° C.) - 1. Color (based on Pt-Co scale) 35 1.1255

EXAMPLE IL

Into a three-neck kettle equipped with a 40 thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Reschig rings and surmounted by a standard refine head with the usual condenser and vacuum receiver attached, were charged 8015 grams (45.0 mols) 45 of refined butyl benzoate and 2414 grams (18.0 mols) of dipropylene glycol. Then 104 grams (1.0 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a pressure 50 of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 175—187° C. The kettle temperature was then reduced slightly, the pressure was reduced to 80 mm. Hg, and the remaining butanol was collected as distillate. A mid-fraction, containing butanol and butyl benzoare was then distilled, followed by a pure fraction of butyl benzoate. Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature below 187° C. The contents of the reaction kettle were cooled, mixed with a filter aid, and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an external nitrogen ebullator. The stripping operation was carried out at a temperature of 210° C. and a pressure of 15 mm. Hg. The kettle residue was then treated with magnesol, filtered and recovered.

A yield of 82.6 per cent of dipropylene glycol dibenzoate (based on dipropylene glycol) was obtained, with an efficiency (based on buryl benzoate) of 87.3 per cent. The refined dipropylene glycol dibenzoate had a purity by saponification of 99.9 per cent, an acidity (as benzoic acid) of 0.06 per cent, a diol content of 0.72 per cent and a color (based on Pt-Co scale) of 35.

EXAMPLE III.

Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Raschig rings and surmounted by a standard reflux head with the usual condenser and vacuum receiver attached, were charged 3565 grams (20.0 mols) of refined butyl benzoate and 1073 grams (8.0 mols) of dipropylene glycol. Then 23 grams (0.5 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 175-180° C. The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg, and the remaining butanol was collected as distillate. A mid-fraction, containing butanol and butyl benzoate was then distilled, followed by a pure fraction of butyl benzoate. Occasional reductions in pressure were required to facilitate 100 maintenance of the kende temperature below 180° C. After most of the excess butyl benzoate had been collected, the pressure was reduced to 0.5 mm. Hg and the temperature of the kettle was raised as rapidly as possible to 105 around 205-207° C. The desired dipropylene glycol dibenzoate was then collected as distillate

A yield of 86.1 per cent of dipropylene glycol dibenzoate (based on dipropylene glycol) was obtained, with an efficiency (based on butyl benzoate) of 78.8 per cent. The refined dipropylene glycol dibenzoate had a purity (by saponification) of 99.9 per cent, an acidity (as benzoic acid) of 0.05 per cent, a diol content of 0.22 per cent and the following physical proporties: properties:-

Refractive index at 20° C. 1.5297 Specific gravity (20/20° C.) -1.1236 Color (based on Gardner scale) Boiling range -

190-195° C. at 0.5 mm. Hg.

EXAMPLE IV.

Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Raschig rings 5 and surmounted by a standard reflux head with the usual condenser and vacuum receiver attached, were charged 3565 grams (20.0 mols) of refined butyl benzoate and 1073 grams (8.0 mols) of dipropylene glycol. Then 4 grams (0.1 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was bested and stirred at a pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle tem-15 perature increased to 175-180° C. The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg, and the remaining butanol was collected as distillate. A mid-fraction, containing butanol and butyl

> Refractive index at 20° C. Specific gravity (20/20° C.) - Color (based on Pt-Co scale) -Boiling range

EXAMPLE V.

40

85

Into a three-neck kettle equipped with a 45 thermowell, a mechanical stirrer and a 25 × 250 mm, column packed with glass Raschig rings and surmounted by a standard reflex bead with the usual condenser and vacuum receiver attached, were charged 1782 grams (4.0 mols) 50 of refined butyl benzoate and 537 grams (10.0 mols) of dipropylene glycol. Then 1.26 grams (0.06 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a pressure of 55 100 mm. Hg. Butanol was removed by distiliation at this pressure until the kertle temperature increased to 175—180° C. The tettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg., and the remaining butanol was collected as distillate. A mid-fraction, containing butanol and butyl benzoare was then distilled; followed by a pure fraction of butyl benzoate. Occasional

> Refractive index at 20° C. Specific gravity (20/20° C) - Color (based on Pt-Co scale) -Boiling range

EXAMPLE VI

Into a three-neck kettle equipped with a 30 thermowell, a mechanical stirrer and a 25 x 250 mm. column packed with glass Raschig rings and surmounted by a standard reflux head with the usual condenser and vacuum receiver attached, were charged 2228 grams (12.5 95 mols) of refined buryl benzoate and 531 grams (5.0 mols) of diethylene glycol. Then 2.76 grams (0.1 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a 100 pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 175° C.

benzoate was then distilled, followed by a pure fraction of buryl benzoate. Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature around 180° C. After most of the excess butyl benzoate had been collected, the pressure was reduced to 2.0 mm. Hg, and the temperature of the kettle was raised as rapidly as possible to around 215—220° C. The desired ester was then collected as distillate.

A yield of 81.2 per cent of dipropylene glycol dibenzoare (based on dipropylene glycol) was obtained, with an efficiency (based on butyl benzoare) of 90.2 per cent. The refined dipropylene glycol dibenzoare had a purity (by saponification) of 99.9 per cent, an acidity (as benzoic acid) of 0.17 per cent, a diol content of 0.19 per cent and the following observed proposition. physical properties:

1,5287

1.1224

210-216° C. at 2 mm. Hg.

reductions in pressure were required to facilitate maintenance of the kettle temperature below 180° C. The contents of the reaction kettle were cooled to 130° C, mixed with a filter aid, and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an 70 external nitrogen chullator. The distillation of dipropylene glycol dibenzoare was carried out at a kettle temperature of 212° C. and a

out at a rettle temperature of 212. C. and a pressure above 1.0 mm. Hg.
A yield of 58.1 per cent of dipropylene glycol dibenzoate (based on dipropylene glycol) was obsained, with an efficiency (based on butyl benzoate) of 74.0 per cent. The refined distance along dibenzoate had a sprint (based on partial distance had a sprin dipropplene glycol dibenzoate had a purity (by appendication) of 98.5 per cent, an acidity (as 80 benzoic acid) of 0.03 per cent, a diol content of 1.50 per cent and the following physical

1.5282 1.1230

35

200-204° C. at 1 mm. Hg.

The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg., and the remaining butanol was collected 105 as distillate. A mid-fraction, containing butanol and butyl benzoate was then distilled followed by a pure fraction of buryl benzoate. Occasional reductions in pressure were required to facilitate maintenance of the kettle 110 temperature below 180° C. The contents of the reaction kettle were conled to 130° C., mixed with a filter sid and filtered to remove The filtrate was then charged the catalyst. to a stripping still equipped with a gooseneck 115 head and an external nitrogen ebullator. distillation of diethylene glycol dibenzoste was

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4 carried out at a temperature of 210-220° C. and a pressure above 1.0 mm. Hg. A yield of 73.7 per cent of diethylene glycol dibenzoate (based on diethylene giycol) was obtained, with an efficiency (based on buryl benzoate) of 76.2 per cent. The refined Refractive index at 20° C. Specific gravity (20/20° C.) - Color (based on Pt-Co scale) -15 Boiling range EXAMPLE VII Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Raschig rings and surmounted by a standard reflex head with the usual condenser and vacuum receiver attached, were charged 2230 grams (12.5 mols) of refined butyl benzoate and 731 grams (5.0 mols) of 2-cthyl-1,3-hexanediol. Then 33 grams (1.0 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 198° (The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg. and the remaining buranol was collected as distillate. A mid-fraction, containing

required to facilitate maintenance of the kettle temperature around 200° C. After most of the excess butyl benzoate had been collected a stripping operation was carried out for ten minutes at a temperature of 220° C, and a pressure of 1.5 mm. Hg. The kettle residue was then filtered and recovered. A yield of 88.4 per cent of 2-ethyl-1,3-

butanol and buryl benzoate was then distilled,

followed by a pure fraction of buryl benzoate.

Occasional reductions in pressure

heranediol dibenzone (based on 2-cityl-1,3hexanediol) was obtained, with an efficiency (based on buryl benzozie) of 84.0 per cent. The refined 2-ethyl-1,3-hexanediol dibenzoate had a purity (by saponification) of 99.4 per cent, an acidity (as benzoic acid) of 0.14 per cent, a diel content of 0.2 per cent and the follow-

ing physical properties:—
Specific gravity (20/20° C.)
Color (based on Gardner scale) 1.0801

Example VIII

Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Raschig rings and surmounted by a standard reflux head with the usual condenser and vacuum receiver attached, were charged 2230 grams (12.5 mols) of refined butyl benzoate and 731 grams (5.0 mols) of 2-ethyl-1,3-hexancdiol. Then 30 grams (10 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a diethylene glycol dibenzoate had a purity (by saponification) of 98.9 per cent, an acidity (as benzoic acid) of 0.07 per cent, a diol content of LO per cent and the following physical properties:-

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1.5438 1.1746 80 200-214° C. at 1 mm. Hg.

pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 196° C. The kettle The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg., and the remaining butanol was collected as distillate. A mid-fraction, containing buranol and buryl benzoate was then distilled, followed by a pure fraction of butyl benzoate. Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature below 200° C. The contents of the reaction kerrie were cooled to around 130° C, mixed with a 80 filter aid and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an external nitrogen ebullator. The stripping operation was carried out at 192° C. and 1 mm. Hg. The kettle residue was then filtered and recovered.

A yield of 75.0 per cent of 2-ethyl-1 hexanediol dibenzoate (based on 2-ethyl-1,3hexanediol) was obtained, with an efficiency (based on butyl benzoate) of 77.5 per cent. The refined 2-ethyl-1,3-hexanediol dibenzoate had a purity (by saponification) of 99.0 per cent, an acidity (as benzoic acid) of 0.18 per cent, a diol content of 0.52 per cent and the following 95 physical properties:-1.0814

Example IX 100

Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 x 250 mm. cclumn packed with glass Raschig rings and surmounted by a standard reflux head with the usual condenser and vacuum receiver 105 attached, were charged 1068 grams (5.99 mols) of refined buryl benzoate and 441 grams (2.5 mols) of 2-methoxymethyl-2,4-dimethyl-15pentanediol. Then 15 grams (1.0 per cent by weight of the kettle charge) of calcium oxide 110 were added and the reaction mixture was heated and stirred at a pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 175--180° C. The kettle temperature was 115 then reduced slightly, the pressure was reduced to 50 mm. Hg., and the remaining butanol was collected as distillate. A mid-fraction, containing butanol and butyl benzoate was then

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distilled, followed by a pure fraction of butyl Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature below 180° C. The con-tents of the reaction kettle were cooled to 130° C., mixed with a filter aid and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an external nitrogen ebullator. Distillation of 2-methoxymethyl-2,4-dimethyl-1,5-pentanediol dibenzoate was carried out at a temperature of 232° C. and a pressure of 1.5 mm. Hg.

25

70

A yield of 83.1 per cent of 2-methoxymethyl-2,4-dimethyl-1,5-pentanediol dibenzoate (based on 2-methorymethyl-2,4-dimethyl-1,5-pentanediol) was obtained, with an efficiency (based on butyl benzoate) of 79.0 per cent. The refined 2 - methoxymethyl-2,4-dimethyl-1,5pennanedial dibenzoate had a purity (by saponification) of 99.8 per cent an acidity (as benzoic acid) of 0.14 per cent, essentially no diol content and the following physical properties:-

Refractive index at 20° C. Specific gravity (20/20° C.) -Color (based on Pr-Co scale) -Boiling point -

1.5310 1.1003 25

225° C. at 1.5 mm. Hg.

EXAMPLE X Into a three-neck kentle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Raschig rings and surmounted by a standard reflex head with the usual condenser and vacuum receiver attached, were charged 446 grams (25 mols) of refined buryl benzoate and 160 grams (1.0 mols) of 2,4-diethyl-1,5-pentanediol. Then 5 grams of calcium oxide were added and the reaction mixture was heated and stimed at a pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 175—189°
C. The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg, and the remaining burand was collected as distillate. A mid-fraction, containing buranol and butyl benzoate was then distilled, folOccasional reductions in pressure required to facilitate maintenance of the kettle temperature below 180° C. The contents of the reaction kettle were cooled to 130° C., mixed with a filter aid and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a guoseneck head and an external nitrogen ebullator. Distillation of 2,4-diethyl-1,5-pentanediol diben-zoate was carried out at a temperature of 254° C and a pressure of 4.0 mm. Hg.
A yield of 36.9 per cent of 2.4-diethyl-1,5-

lowed by a pure fraction of butyl benzoate. Refractive index at 20° C. Specific gravity (20/20° C) - Color (based on Pt-Co scale) -Boiling point

pentanedial dibenzoate (based on 2.4-diethyl-1,5-pentanediol) was obtained, with an efficicary (based on buryl benzoate) of 44.8 per cent. The refined 2,4-diethyl-1,5-pentane-diel dibenzoate had a purity (by saponification) of 99.6 per cent, an acidity (as benzolc acid) of 0.62 per cent, a diol content of 0.04 per cent and the following physical properties:-

EXAMPLE XI

1.5322 1.0775

206° C, at 1 mm, Hg,

Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Raschig rings and surmounted by a standard reflux head with attached, were charged 8015 grams (45 mols) 80 of refined butyl benzoate and 2125 grams (18 mole) of 3-methyl-1.5-pentanediol. Then 101 the usual condenser and vacuum receiver mole) of 3-methyl-1,5-pentanediol. Then 101 grams (1.0 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 190-200° The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm.
Hg., and the remaining but mol was collected as distillate. A mid-fraction, containing but anol and butyl benzoate was then distilled, followed by a pure araction of butyl benzoate.

Occasional reductions in pressure required to facilitate maintenance of the bettle temperature below 190° C. The contents of the reaction bettle were cooled, mixed with a filter aid and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an exter- 100 nal nitrogen ebullator. The stripping operation was carried out at a kettle temperature of 180-189° C, and a pressure of 2 to 4 mm. Hg. The kettle residue was then filtered and

A yield of 85.3 per cent of 3-methyl-1,5pentanediol dibenzoate (based on 3-methyl-1,5pentanedici) was obtained, with an efficiency (based on buryl benzoare) of 86.6 per cent. The refined 3-methyl-1,5-pentanedici dibenzoate had a purity (by saponification) of 100.2 per cent, an acidity (as benzoic acid) of 0.01 per cent, essentially no diol content and the following physical properties:

50

95

Refractive index at 20° C. - - 1.5378

Specific gravity (20/20° C.) - - 1.1103

Color (based on Pt-Co scale) - 70

Boiling range - - - 186—192° C. at 1 mm. Hg.

Freezing point - - - - - - 33.4° C.

EXAMPLE XIF

Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25×250 mm. column packed with stainless steel promm. column packed with stainless steel protection traded packing and surmounted by a standard reflux head with the usual condenser and vacuum receiver attached, were charged 1338 grams (7.5 mols) of refined butyl benzoate and 480 grams (3.0 mols) of 2-cthyl-2-butyl-1,3-15 propanediol. Then 18.2 grams (1.0 per cent by weight of the kettle charge) of calcium oxide were added and the reaction mixture was heated and stirred at a pressure of 90 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature was then reduced slightly, the pressure was reduced to 75 mm. Hg., and the remaining butanol was collected as distillate. A midfraction, containing butanol and butyl benzoate was then distilled, followed by a pure fraction of butyl benzoate. Occasional reduced.

Refractive index at 20° C. Specific gravity (20/20° C.) - Color (based on Pt-Co scale) - Boiling range - - -

EYAMPLE XIII

Into a three-neck kettle equipped with a 55 thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with stainless steel protruded packing and surmounted by a standard reflux head with the usual condenser and vacuum receiver attached, were charged 200 grams (1.5 mols) of 2-methyl-2-propyl-1,3-propenedial and 670 grams (3.75 mals) of refined butyl benzoate. Then 9 grams of calcium oxide (1.0 per cent by weight of the kettle charge) were added and the reaction mixture was heated and stiered at a pressure of 100 mm. Hg. Buranol was removed by distilla-tion at this pressure until the kettle tempera-ture rose to 156° C. The kettle temperature was reduced slightly, the pressure was reduced was reduced slightly, the pressure was reduced to 50 mm. Hg., and the remaining butanol was collected as distillate. A mid-fraction, containing butanol and butyl benzoate was then

Refractive index at 20° C. Specific gravity (20/20° C.) - Color (based on Pt-Co scale) - Boiling range - - - - Freezing point - - - -

EXAMPLE XIV

In a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with stainless steel protruded saddles, and surmounted by a standard reflux head with the usual condenser and

tions in pressure were required to facilitate maintenance of the kertle temperature below 160° C. The contents of the reaction kertle were cooled, mixed with a filter aid and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an external nitrogen chullator. Distillation of 2-ethyl-2-butyl-1,3-propanediol dibenzoate was carried out at a temperature of 237—267° C. and a pressure

of 1.5 mm. Hg.

A yield of 64.7 per cent of 2-ethyl-2-butyl-1,3-propanediol dibenzoate (based on 2-ethyl-2-butyl-1,3-propanediol) was obtained, with an efficiency (based on butyl benzoate) of 87.4 per cent. The refined 2-ethyl-2-butyl-1,3-propanediol dibenzoate had a purity (by saponification) of 100.8 per cent, an acidity (as benzoic acid) of 0.02 per cent, diol content of 0.39 per cent and the following physical properties:—

1.5327 1.0778 20

236° C. at 1.5 mm. Hg.

distilled, followed by a pure fraction of buryl benzoate. Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature below 170° C. The contents of the reaction kettle were cooled, mixed with a filter sid and filtered to remove the canalyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an external nitrogen ebullator. Distillation of the ester was carried out at a kettle temperature of 249—252° C. and a pressure of 2.0

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mm. Hg.

A yield of 48.2 per cent of 2-methyl-2-propyl-1,3-propanediol dibenzoate (based on 2-methyl-2-propyl-1,3-propanediol) was obtained. The refined 2-methyl-2-propyl-1,3-propanediol dibenzoate had a purity of 101.8 (by saponification), essentially no acidity and diol content and the following physical proper-

- 1.5367 - 1.1003 - 20 - 214° C. at 2 mm. Hg. - —5.6° C.

vacuum receiver attached, were charged 670 grams (3.75 mols) of refined buryl benzoate and 200 grams (1.5 mols) of 2,2-diethyl-1,3-propanediol. Then 9 grams of calcium oxide (1.0 per cent by weight of the kettle charge) were added and the reaction mixture was

heated and stirred at a pressure of 100 mm. Butanol was removed by distillation at this pressure until the kettle temperature increased to 165° C. The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg., and the remaining butanol was collected as distillate. A midfraction containing butanol and butyl benzo-ate was then distilled, followed by a pure fraction of buryl benzoate. Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature below 184° C. The contents of the reaction kettle were cooled, mixed with a filter aid and filtered to remove the catalyst. The filtrate was then Refractive index at 20° C.

Specific gravity (20/20° C.) - Color (based on Pt-Co scale) -Boiling range Freezing point -

Example XV

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Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with glass Raschig rings and surmounted by a standard reflux head with the usual condenser and vacuum receiver attached, were charged 802 grams (4.5 mols) of refined butyl benzoate and 236 grams (2.0 mols) of 2-methyl-1,5-pentanediol. Then 10 grams of calcium oxide (1.0 per cent by weight of the kettle charge) were added and the reaction mixture was heated and stirred at a pressure of 100 mm. Hg. Butanol was removed by distillation at this pressure until the kettle temperature increased to 171° C. The kettle temperature was reduced slightly, the pressure was reduced to 50 mm. Hg., and the remaining butanol was collected as distillate. A mid-fraction, containing butanol and butyl benzoate was then distilled, followed by a pure

Refractive index at 20° C. Specific gravity (20/20° C.) - Color (based on Pt-Co scale) -Boiling range -Freezing point -

EXAMPLE XVI

Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250 mm. column packed with stainless steel protruded packing and surmounted by a standard 85 reflux head with the usual condenser and vacuum receiver attached, were charged 850 grams (4.77 mols) of refined butyl benzoate and 264 grams (2.0 mols) of 2-ethyl-1,5-pen-tanediol. Then 11 grams of calcium oxide (1.0 per cent by weight of the kettle charge) added and the reaction mixture was heated and stirred at a pressure of 100 mm. Butanol was removed by distillation at this pressure until the kettle temperature increased to 154° C. The kettle temperature was then reduced slightly, the pressure was reduced to 50 mm. Hg., and the remaining charged to a stripping still equipped with a gooseneck head and an external nitrogen ebullator. Distillation of the ester was carried out at a kettle temperature of 200° C. and a pressure of 2.0 mm. Hg.

A yield of 42.9 per cent of 2,2-diethyl-1,3propanediol dibenzoste (based on 2,2-diethyl-1,3-propandial) was obtained, with an efficiency (based on butyl benzoate) of 51.1 per cent. The refined 2,2-diethyl-1,3-propanediol dibenzoate had a purity of 101.2 per cent (by seponification), essentially no acidity, a diol content of 0.34 per cent, and the following physical properties:-

1.5402 1.1097 18 211-

-214° C, at 2 mm. Hg. -5° C.

fraction of butyl benzoate. Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature below The contents of the reactor kettle were cooled, mixed with a filter aid and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an external mirrogen chul-Distillation of the ester was carried lator. out at a kettle temperature of 200 to 210° C.

at 2.0 mm. Hg.
A yield of 88.3 per cent of 2-methyl-1, pentanedial dibenzoate (based on 2-methyl-1,5-pentanedial) was obtained, with an efficiency of 88.8 (based on butyl benzoate). The refined 2-methyl-1,5-pentanediol dibenzoure had a purity of 99.8 per cent (by saponification) an acidity (as bearoic acid) of 0.12, essentially no diol content and the following physical properties:

1.5379 1.1045 18 -204° C, at 2 mm. Hg. 203--3.8° C.

butanol was collected as distillate. fraction, containing butanel and butyl benzo-ate was then distilled, followed by a pure fraction of butyl benzoate, Occasional reductions in pressure were required to facilitate maintenance of the kettle temperature below 187° C. The contents of the reaction kettle were cooled, mixed with a filter aid and filtered 105 to remove the caralyst. The filtrate was then charged to a stripping still equipped with a gooseneck head and an external nirrogen chul-Distillation of the ester was carried our at a kettle temperature of 218-221° C. 110

A yield of 89.8 per cent of 2-ethyl-1,5-pen-tanediol dibenzoate (hased on 2-ethyl-1,5-pen-tanediol) was obtained with an efficiency (based on butyl benzoate) of 91.4 per cent.

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	refined 2-ethyl-1,5-pentanedial dibenzoate had a punity (by seponification) of 99.9 per cent, an acidity (as benzoic acid) of 0.13 per cent,	and a diol content of 0.69 per cent, and the following physical properties:—	5
10	Refractive index at 20° C Specific gravity (20/20° C.) - Color (based on Pr-Co scale) - Boiling range Freezing point	- 1.5356 - 1.0945 - 20 - 215—218° C, at 1.5 mm. Hg. - 35° C.	
	EXAMPLE XVII Into a three-neck kettle equipped with a thermowell, a mechanical stirrer and a 25 × 250	butyl benzoate. Occasional reductions in pressure was required to facilitate mainten-	
15	nm. column packed with glass Raschig rings and summounted by a standard refirst head with the usual condenser and vacuum receiver attached, were charged 620 grams (3.48 mols)	ance of the kettle temperature below 212° C. The contents of the reaction kettle were cooled, mixed with a filter aid and filtered to remove the catalyst. The filtrate was then charged to a stripping still equipped with a gooseneck	35
20	of refined buryl benzoare and 200 grams (1.31 mols) of 2-ethyl-3-methyl-1,5-pentanediol. Then 8 grams (1.0 per cent by weight of the kettle charge) of lime (technical grade calcium	head and an external nitrogen ebullator. Dis- tillation of the ester was carried out at a kettle temperature of 217—222° C. and a pressure of 2.0 mm, Hg.	40
25	oxide) were added and the reaction mixture was heated to 110° C, and stirred at a pressure of 100 mm, Hg. Butanol was removed by	A yield of 860 per cent of 2-ethyl-3-methyl-1,5-pentanedial dibenzoate (based on 2-ethyl-3-methyl-1,5-pentanedial) was obtained, with an efficiency (based on butyl benzoate) of 79,7	45
	perature increased to 190° C. The kettle temperature was then reduced slightly, the pressure was reduced, and the remaining buranol was collected as distillate. A mid-fraction,	per cent. The refined 2-ethyl-3-methyl-1,5- penumedial dibenzane had a purity of 98.7 per cent (by saponification), an acidity (as benzoic	
30	containing butanol and butyl benzoate was then distilled, followed by a pure fraction of	acid) of 0.12 per cent, a diol content of 0.41 per cent and the following physical properties:—	50
55	Refractive index at 20° C Specific gravity (20/20° C.) - Color (based on Pt-Os scale) - Boiling range Freezing point	- 1.5362 - 1.0945 - 18 - 210—218° C. at 2 mm, Hg. 17.2° C.	
60	WHAT WE CLAIM IS:— 1. Process for preparing dibenzoate esters of aliphatic diods which comprises reacting butyl benzoate with an aliphatic diod contain-	of from 2 mm. of mercury to 100 mm. of mer- cury. 9. Process as claimed in any of Claims 1 to 8 in which the aliphatic diol is dipropylene	85
	ing 4 to 9 carbon atoms and having no terriary hydroxyl groups as hereinhefore defined. 2. Process as claimed in Claim 1 in which	giycol. 10. The dibenzoate ester of 2-ethyl-1,3-hexane diol.	90
65	one molecular proportion of the aliphatic diol is reacted with 25 molecular proportions of butyl benzoate. 3. A process as claimed in Claim 1 or 2 in	11. The dibenzoate ester of 2-methyl-1,5-pentanediol. 12. The dibenzoate ester of 2-ethyl-1,5-pentanediol.	
70	which the reaction is effected in the presence of an alkaline catalyst. 4. Process as claimed in Claim 3 in which	pentanediol. 13. The dibenzoate ester of 2-ethyl-3-methyl-1,5-pentanediol. 14. The dibenzoate ester of 2,4-diethyl-1,5-	95
75	the alkaline caralyst is an alkali metal or alka- line earth metal oxide, hydroxide, alkoxide, car- bonate or borate. 5. Process as claimed in Claim 4 in which	pentanediol. 15. The dibenzoate ester of 2-cthyl-2-butyl-13-propanodiol.	100
	the alkaline earth metal oxide is calcium oxide. 6. Process as claimed in Claim 3, 4 or 5 in which the catalyst concentration is between	16. The dibenzoate ester of 2-methyl-2-propyl-1,3-propanediol. 17. Process for preparing dibenzoate esters of aliphatic diols substantially as herein des-	105
80	0.1% and 1.0% by weight. 7. Process as claimed in any of Claims 1 to 6 in which the reaction is effected at a tem-	cribed with reference to and as illustrated in the foregoing examples.	
	perature of from 150° C. to 180° C. 8. Process as claimed in any of Claims 1 to 7 in which the reaction is effected at a pressure	W. P. THOMPSON & CO., 12, Church Street, Liverpool, 1, Chartered Patent Agents.	

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